Diminished Potential for B-Lymphoid Differentiation after Murine Leukemia Virus Infection In Vivo and in EML Hematopoietic Progenitor Cells

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Infection with a recombinant murine-feline gammaretrovirus, MoFe2, or with the parent virus, Moloney murine leukemia virus, caused significant reduction in B-lymphoid differentiation of bone marrow at 2 to 8 weeks postinfection. The suppression was selective, in that myeloid potential was significantly increased by infection. Analysis of cell surface markers and immunoglobulin H gene rearrangements in an in vitro model demonstrated normal B-lymphoid differentiation after infection but significantly reduced viability of differentiating cells. This reduction in viability may confer a selective advantage on undifferentiated lymphoid progenitors in the bone marrow of gammaretrovirus-infected animals and thereby contribute to the establishment of a premalignant state.

Feline leukemia virus (FeLV) and murine leukemia virus (MuLV) are closely related gammaretroviruses that induce malignant tumors of hematopoietic and lymphoid tissues through a complex, multistep process. Previous studies have demonstrated the significant impact of gammaretrovirus infection on bone marrow hematopoiesis in the early stages of disease induction (1, 8, 11–13, 21, 28). For example, Moloney MuLV (M-MuLV) infection induces compensatory extramedullary hematopoiesis, a result of diminished support of hematopoiesis by bone marrow stromal cells (2, 6, 8, 24). Altered progenitor cell distribution has also been observed at 2 to 4 weeks postinfection with SL3-3 MuLV (21). Virus replication in the bone marrow, and consequent disruption of hematopoiesis, is also a hallmark of FeLV infection (1, 13, 14, 18, 20). In a recent longitudinal study, significant depression in circulating red blood cell counts and segmented neutrophils was observed in infected cats during the first 4 weeks postinoculation (4).

FeLV-945 is a unique isolate of FeLV naturally associated with malignant, degenerative, and proliferative disorders of non-T-cell origin. Experimental infection with a virus bearing the unique genetic hallmarks of FeLV-945 resulted in the rapid induction of a multicentric lymphoma of B-lymphoid origin, in contrast to the long-latency T-cell lymphomas characteristic of natural, horizontally transmissible FeLV (3, 4). The basis for the shift in tumor spectrum remains unknown; however, the long terminal repeat (LTR) element of FeLV-945 contains a unique repeat motif comprised of a 21-bp element triplicated in tandem beginning 25 bp downstream of the canonical transcriptional enhancer (5). To investigate the influence of the FeLV-945 LTR on bone marrow hematopoiesis, a murine-feline recombinant retrovirus was constructed by substituting the triplicate-containing U3 region of FeLV-945 for that of M-MuLV. The recombinant virus, termed MoFe2-MuLV (MoFe2), infects mice and induces T-cell lymphomas with kinetics comparable to those of M-MuLV (10, 23).

To determine whether MoFe2 infection causes extramedullary hematopoiesis and splenomegaly early in disease progression, neonatal NIH/Swiss mice were inoculated intraperitoneally with equivalent amounts of MoFe2 or M-MuLV as normalized by reverse transcriptase activity. Groups of four to six animals were sacrificed at regular intervals 2 to 8 weeks postinoculation, and spleen weight was determined as a percentage of body weight. The results demonstrated significantly greater spleen weights in M-MuLV-infected mice than in MoFe2-infected or mock-infected animals at 2, 4, 6, and 8 weeks postinoculation (Fig. 1A). Pre-B clonogenic progenitor cells were quantified by depositing unfractionated bone marrow from individual animals was deposited at densities of 2 × 10³ nucleated cells per ml in triplicate into medium containing 50 ng/ml stem cell factor, 10 ng/ml interleukin 3 (IL-3), 10 ng/ml IL-6, and 3 units/ml erythropoietin in methylcellulose (MethoCult M3434; StemCell Technologies, Inc., Vancouver, Canada). Significantly increased CFU–granulocyte-erythrocyte-monocyte-megakaryocyte (GEMM) clonogenic potential was recovered from the bone marrow of animals infected with either virus at 6 and 8 weeks postinfection; increased CFU–granulocyte-monocyte/macrophage (GM) clonogenic potential was also detected in the bone marrow of infected animals at 8 weeks postinfection (Fig. 1A). Pre-B clonogenic progenitor cells were quantified by depositing unfracti...
FIG. 1. Colony-forming assays of bone marrow and EML cells following infection with MoFe2 or M-MuLV. (A) Unfractionated bone marrow collected at intervals after mock infection or infection with MoFe2 or M-MuLV was deposited in semisolid culture medium optimized for the differentiation and growth of myeloid colony-forming progenitors (CFU-GEMM, CFU-GM) or pre-B-cell colony-forming progenitors (CFU-pre-B). After incubation for 14 days (myeloid) or 7 days (pre-B lymphoid), colonies were typed and enumerated by light microscopy. Statistical analysis using one-way ANOVA showed significantly increased CFU-GEMM ($P < 0.05$) at 6 and 8 weeks postinfection with either virus, significantly increased CFU-GM ($P < 0.05$) at 8 weeks postinfection with either virus, significantly decreased CFU-pre-B ($P < 0.05$) at 2, 4, and 6 weeks postinfection with MoFe2, and significantly decreased CFU-pre-B ($P < 0.05$) at 8 weeks postinfection with M-MuLV compared to age-matched mock-infected control animals. (B) Infected and uninfected EML cells were stimulated with $10^{-5}$ M all-trans retinoic acid and 10 ng/ml IL-3 to remove the block to myeloid differentiation and were then plated in semisolid culture medium optimized for the differentiation and growth of myeloid colony-forming progenitors (CFU-GEMM). Undifferentiated blast colonies form under these conditions as well (CFU-Blast). EML cells were also plated in medium optimized for the differentiation and growth of pre-B-cell colony-forming progenitors (CFU-pre-B). After incubation for 14 days, colonies were typed and enumerated by light microscopy. Statistical analysis using one-way ANOVA showed significantly decreased CFU-Blast ($P < 0.05$), significantly increased CFU-GEMM ($P < 0.05$), and significantly decreased CFU-pre-B ($P < 0.05$) after infection with either virus.
FIG. 2. Temporal expression of the surface markers CD45R/B220, CD19, and CD43 on EML cells stimulated to undergo B-lymphoid differentiation. Expression of surface markers on MoFe2-infected and M-MuLV-infected EML cells was examined by flow cytometry using phycoerythrin-conjugated monoclonal antibodies (BD Biosciences, Palo Alto, CA). Included as controls were uninfected EML cells stimulated in parallel to undergo B-lymphoid differentiation (Uninf.) and uninfected EML cells in the absence of stimulation (Uninf. unstimulated). Uninfected, unstimulated cells examined with an isotype control antibody were included as a negative control. Histograms depict 10,000 events per sample, and results are representative of three independent experiments. The results shown in panel A represent day 6 of differentiation. The results shown in panel B represent day 6, 12, or 15 of differentiation, as indicated.
bone marrow from individual animals at a density of $5 \times 10^5$ nucleated cells per ml in triplicate into medium containing 10 ng/ml IL-7 in methylcellulose (MethoCult M3630; StemCell Technologies, Inc., Vancouver, Canada). In contrast to the myeloid hyperplasia, a significant decrease in CFU–pre-B-cell clonogenic potential was observed at 2, 4, and 6 weeks postinfection for MoFe2-infected mice and at 8 weeks postinfection for M-MuLV-infected mice (Fig. 1A). This observation represents the first report of suppression of the B-lymphoid differentiation potential in bone marrow from gammaretrovirus-infected animals during the preleukemic stage.

Studies of the impact of gammaretrovirus infection on hematopoiesis have generally analyzed bone marrow removed directly from the animal or from long-term bone marrow cultures, both of which are complex, multicomponent systems. To simplify this analysis, we used the murine lymphohematopoietic progenitor cell line EML, a bone marrow-derived, stem cell factor-dependent progenitor cell line capable of differentiation along erythroid, myeloid, and lymphoid lineages. EML was established from normal mouse bone marrow by transducing a retrovirus vector that expresses a dominant-negative retinoic acid receptor (26). Previous reports reveal that EML cells represent a reliable and authentic model of hematopoietic progenitor lineage commitment and differentiation (7, 15, 19, 27).

To examine myeloid clonogenic potential, EML cells were stimulated with $10^{-5}$ M all-trans retinoic acid (Sigma, St. Louis, MO) and 10 ng/ml IL-3 (Peprotech, Rocky Hill, NJ) for 72 h and were then deposited in triplicate at $10^5$ cells per ml into MethoCult M3434 medium. After 14 days of incubation, colonies were typed and enumerated by light microscopy. A significantly decreased number of undifferentiated CFU-blast colonies were typed and enumerated by light microscopy after 7 days of incubation. Consistent with observations of infected bone marrow, EML cells demonstrated significantly decreased CFU–pre-B-cell clonogenic potential after infection with either MoFe2 or M-MuLV (Fig. 1B).

To examine the stage of B-lymphoid differentiation affected by infection, cells were cultured in Iscove’s modified Dulbecco medium (Invitrogen, Carlsbad, CA) supplemented with 20% horse serum, 50 ng/ml IL-7, and 10 ng/ml BMP-2, refreshed every 2 to 3 days. B-lymphoid differentiation was monitored by flow cytometry to measure expression of the temporally regulated cell surface markers CD45R/B220, CD43, CD19, and CD25. As described by others, CD45R/B220 is expressed early and throughout differentiation; CD43 is expressed early, persists through immunoglobulin H (IgH) D-J gene rearrangement, and declines at the pro-B-cell stage; CD19 appears at the time of IgH D-J gene arrangement and persists; and CD25 is not expressed until the time of IgH VDJ gene rearrangement at the pre-B cell stage (9, 16, 17, 22). In our study, CD45R/B220 was detectable by day 3 and persisted throughout the study equally in uninfected and infected cells (Fig. 2A).

To examine the clonogenic potential of infected bone marrow, single cells were placed directly into amplification reactions by infection, cells were cultured in Iscove's modified Dulbecco medium supplemented with 20% horse serum, 50 ng/ml IL-7, and 10 ng/ml BMP-2, refreshed every 2 to 3 days. B-lymphoid differentiation was monitored by flow cytometry to measure expression of the temporally regulated cell surface markers CD45R/B220, CD43, CD19, and CD25. As described by others, CD45R/B220 is expressed early and throughout differentiation; CD43 is expressed early, persists through immunoglobulin H (IgH) D-J gene rearrangement, and declines at the pro-B-cell stage; CD19 appears at the time of IgH D-J gene arrangement and persists; and CD25 is not expressed until the time of IgH VDJ gene rearrangement at the pre-B cell stage (9, 16, 17, 22). In our study, CD45R/B220 was detectable by day 3 and persisted throughout the study equally in uninfected and infected cells (Fig. 2A).

**Table 1. Ig heavy-chain gene rearrangement in infected and uninfected EML cells as measured by single-cell PCR**

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<tr>
<th>EML cell sample</th>
<th>% Heavy-chain gene rearrangement</th>
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<tr>
<td></td>
<td>Germ line</td>
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<tr>
<td></td>
<td>Uninfected</td>
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<td>100</td>
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*Uninfected EML cells or cells infected with MoFe2 or M-MuLV were stimulated to B-lymphoid differentiation and collected at regular intervals for 15 days. At each collection, single cells were deposited directly into amplification reactions by limiting dilution. Single-cell PCR amplification was performed with multiple primer sets to detect the germ line IgH J region, D-J rearrangements, and VDJ rearrangements involving three of the largest VH families (J558, Q52, and 7183) by a procedure described by others (25). Eight independent amplifications were examined with each primer set at each time point, and the results are reported as percentages demonstrating the indicated target, ND, not done.

Rearrangement of the Ig heavy-chain gene (IgH) was then examined with a single-cell PCR procedure described by others (25) and recently modified in a manner to be detailed elsewhere (M. Gunthart and N. Rosenberg, unpublished data). Infected and uninfected EML cells were collected at regular intervals for 15 days during B-lymphoid differentiation, and single cells were placed directly into amplification reactions by limiting dilution. Samples were amplified in two sequential rounds of PCR using multiple primer sets designed to amplify the germ line IgH J region, D-J rearrangements, and VDJ rearrangements involving three of the largest VH families. Loss of germ line configuration was detected between 3 and 6 days in all populations. D-J rearrangements were first apparent by day 3 and were evident at comparable frequencies in all populations thereafter (Table 1).

Because a delay or block in differentiation did not account for the reduction in the B-lymphoid potential of infected EML cells, we examined effects on cell viability. EML cells were deposited at $5 \times 10^5$ cells per ml and stimulated to undergo B-lymphoid differentiation. Cell viability was mea-
FIG. 3. Diminished viability of infected EML cells after stimulation to B-lymphoid differentiation. EML cells infected with MoFe2 or M-MuLV and uninfected control cells were stimulated to undergo B-lymphoid differentiation as described in the text. Cell viability was measured at 3 days poststimulation by MTS dye reduction assay (Promega, Madison, WI). Statistical analysis by one-way ANOVA showed significantly decreased viability (P < 0.05) of EML cells infected with either virus. The data shown are averages (± standard errors) from triplicate experiments.


25. ten Boekel, E., F. Melchert, and A. Rolink. 1995. The status of Ig loci
rearrangements in single cells from different stages of B cell development. 